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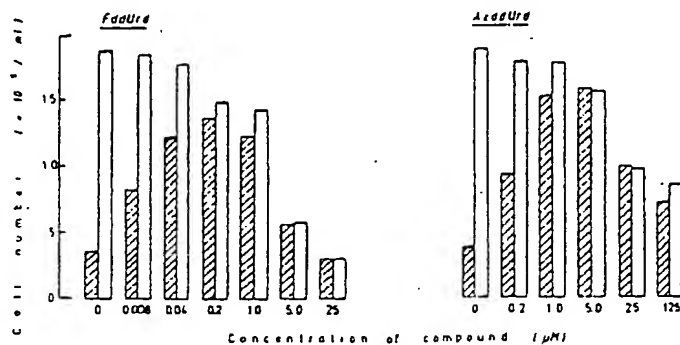
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(54) 3'-Fluoro-2',3'-dideoxyuridine, and its therapeutic application.

(57) The novel compound 3'-fluoro-2',3'-dideoxyuridine has an antiretroviral effect against human immunodeficiency virus (HIV) and is a potential antiretroviral agent for treating AIDS and related viral diseases, including hepatitis B.

FIG 2



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3'-Fluoro-2',3'-dideoxyuridine, and its therapeutic application

This invention relates to a novel 3'-fluoro-pyrimidine-2',3'-dideoxynucleoside, 3'-fluoro-2',3'-dideoxyuridine, and its application as a novel therapeutic agent for treating retroviral diseases, including hepatitis B, and for treating AIDS and AIDS-related diseases.

Since the identification of a retrovirus, referred to as human immunodeficiency virus (HIV), as the causative agent of the acquired immunodeficiency syndrome (AIDS), many attempts have been made to develop agents that may be useful in the prevention and/or therapy of this disease. Among the 2',3'-dideoxynucleosides, several congeners have been reported to show promising antiretroviral activity. It is demonstrated that 3'-azido-2',3'-dideoxythymidine (AzddThd) is a potent inhibitor of HIV replication and protects ATH8 lymphocytes against the cytopathic effect of the virus at a concentration of 5 to 10 μ M. Moreover, AzddThd protects MT4 lymphocytes against HIV at a concentration as low as 0.02 μ M. 2',3'-didehydro-2',3'-dideoxythymidine and 2',3'-dideoxythymidine completely protect MT4 cells against HIV at 0.04 μ M and 1 μ M respectively. 2',3'-dideoxycytidine (ddCyd) is a potent inhibitor of HIV-induced cytopathogenicity in ATH8 cells, and it was shown that 2',3'-didehydro-2',3'-dideoxycytidine (ddeCyd) had a similar antiretroviral activity as ddCyd in ATH8 cells. This observation is later confirmed for MT4 cells and for human peripheral mononuclear blood cells. Interestingly, among the purine-2',3'-dideoxynucleosides, 2',3'-dideoxyadenosine (ddAdo), 2',3'-dideoxyinosine (ddIno) and 2',3'-dideoxyguanosine (ddGuo) are equally effective in protecting ATH8 cells against the cytopathogenicity of HIV (Mitsuya, H., et al. Proc. Natl. Acad. Sci. USA 83, 1911-1915 (1986)). Moreover, in ATH8 cells, the selectivity index (the ratio of the cytotoxic dose of the effective dose) was higher for ddAdo than for any of the other purine-2',3'-dideoxyriboside tested (Mitsuya, H., et al. Broder, S., Ed., Marcel Dekker Inc., New York, pp 303-333, (1986)).

During extensive research for novel potential inhibitors of HIV, a novel 3'-fluoro-substituted pyrimidine-2',3'-dideoxynucleoside was found, namely 3'-fluoro-2',3'-dideoxyuridine (FddUrd). We found that the novel FddUrd derivative is 10-fold more potent against HIV than AzddUrd.

Hereafter the anti-retrovirus potency of FddUrd will be compared with other 3'-fluoro- and 3'-azido-substituted derivatives of 2',3'-dideoxyuridine (AzddUrd), 2',3'-dideoxy-5-ethyl-uridine (FddEtUrd, AzddEtUrd) and 2',3'-dideoxycytidine (FddCyd, AzddCyd).

All reagents used were of the highest quality obtainable.

2',3'-Dideoxycytidine (ddCyd) was obtained from Pharmacia PL-Biochemicals (Piscataway, NJ, USA). 3'-Azido-2',3'-dideoxycytidine was a kind of gift of Dr. D.G. Johns (National Cancer Institute (NCI), National Institutes of Health (NIH), Bethesda, MD, USA). The method of J.P. Horwitz et al (J. Org. Chem. 31, 205-211 (1966)) was used to synthesize 2',3'-dideoxyuridine (ddUrd). 2',3'-Dideoxy-5-ethyluridine (ddEtUrd), 3'-azido-2',3'-dideoxy-5-ethyluridine (AzddEtUrd), 3'-fluoro-2',3'-dideoxy-5-ethyluridine (FddEtUrd), and 3'-fluoro-2',3'-dideoxycytidine (FddCyd) were synthesized as described by P. Herdewijn et al (J. Med. Chem. 30, 1270-1278 (1987)). 3'-fluoro-2',3'-dideoxyuridine has been synthesized starting from 1-(5-O-trityl-2-deoxy- β -D-threo-pentofuranosyl)uracil (Lin T.-S., et al, J. Med. Chem. 26, 544, (1983)) with diethylaminosulfur trifluoride, followed by deprotection of the 5'-O-trityl group with 80% of acetic acid at 100°C. The synthesis of 3'-azido-2',3'-dideoxyuridine (AzddUrd) is described by R.F. Schinazi et al (Abstracts of the Twenty Seventh Interscience Conference on Antimicrobial Agents and Chemotherapy, New York, 4-7 October 1987, p. 161, no. 369 (1987)).

The various types of cells used in the experiments below were grown as follows.

Raji/O cells and Raji/TK⁻ (a thymidine (dThd) kinasedeficient mutant cell line derived from wild-type Raji/O cells, were grown in 75 cm² plastic tissue culture flasks (International Medical, Brussels, Belgium), in Eagle's minimal essential medium, supplemented with 10% (v/v) inactivated foetal calf serum (Gibco, Glasgow, Scotland, U.K.), 2 mM L-glutamine (Flow Laboratories, Irvine, Scotland, U.K.) and 1% THM (5 mM Tes, 7.5 mM Hepes and 5 mM (MOPS) buffer (pH 5.3). The Raji/O and Raji/TK⁻ cells were characterized according to J. Balzarini et al (Biochem. Pharmacol. 31, 1089-1095 (1982)). Molt/4F, CEM and H9 cells were grown in Eagle's minimal essential medium, supplemented with 10% (v/v) inactivated foetal calf serum and 2 mM L-glutamine. MT4 cells were cultivated in RPMI 1640 medium (Gibco) containing/20 mM Hepes buffer, 10% (v/v) inactivated foetal calf serum (Gibco) and 2 mM L-glutamine (Flow Laboratories).

The two types of viruses used in the experiments below, were prepared from distinct sources as follows.

HTLV-III_B (designated HIV) was derived from a pool of American patients with AIDS, and obtained from the supernatant of HIV-infected H9 cell cultures (see M. Popovic et al, Science 224, 497-500 (1984)).

Moloney murine sarcoma virus (MSV) was prepared from tumors induced by *in vivo* infection of 10 day-old NMRI mice according to the procedure described by De Clercq et al (Proc. Soc. Exp. Biol. Med. 137, 590-594 (1971)).

Therapeutic compositions containing 3'-fluoro-2',3'-dideoxyuridine as an active ingredient for treating retroviral diseases, in particular AIDS or AIDS-related diseases, including hepatitis B in human practice, may take the form of powders, suspensions, solutions, sprays, emulsions, unguents or creams and may be used for local application, for intranasal, rectal, vaginal and also for oral or parenteral (intravenous, intradermal, intramuscular, intrathecal etc.) administration. Such compositions may be prepared by combining (e.g. mixing, dissolving etc.) the active compound of 3'-fluoro-2',3'-dideoxyuridine with pharmaceutically acceptable excipients of neutral character (such as aqueous or non-aqueous solvents, stabilizers, emulsifiers, detergents, additives), and further, if necessary with dyes and aromatizers. The concentration of the active ingredient in the therapeutic composition may vary widely between 0.1% and 100%, dependent on the mode of administration. Further, the dose of the active ingredient to be administered may vary between 0.1 mg and 100 mg per kg of body weight.

The pharmaceutical properties of FddUrd, notably its anti-retroviral action, especially against HIV, are documented in comparison with other 3'-fluoro- and 3'-azido-substituted pyrimidine-2',3'-dideoxynucleosides, by the following examples which should not be read in a restrictive sense.

In the examples reference is made to the drawings annexed, in which:

Fig. 1 shows the structural formulae of 3'-fluoro- and 3'-azido-substituted pyrimidine 2',3'-dideoxynucleosides, tested;

Fig. 2 is a representation of the test results on the inhibition of the cytopathogenicity of HIV for MT4 cells by FddUrd and AzddUrd. Viability of the cells was measured by trypan blue exclusion after an incubation period of 5 days. Mock-infected cells, incubated in the presence of different concentrations of the test compounds are indicated by open columns (□); HIV-infected cells, incubated in the presence of different concentrations of the test compounds are indicated by black columns (■). Data represent average values for 3 to 5 separate experiments;

Fig. 3 shows double-reciprocal plots for inhibition of Mol/4F dThd kinase by FddUrd (A), and AzddUrd (B). Inhibitor concentrations: none (●), 1000 μ M (Δ) and 2000 μ M (□) for FddUrd; none (●), 500 μ M (Δ) and 1000 μ M (□) for AzddUrd. Data represent average values for 3 separate experiments; and

Fig. 4 shows double-reciprocal plots for inhibition of Mol/4F dCyd kinase by FddCyd (A) and AzddCyd (B). Inhibitor concentrations: none (●), 1000 μ M (□) and 200 μ M (Δ) for FddCyd; none (●) and 250 μ M (Δ) for AzddCyd. Data represent average values for 3 separate experiments.

EXAMPLE 1

Anti-retrovirus effects of FddUrd

FddUrd, the unsubstituted analogue and other 3'-fluoro- and 3'-azido-substituted analogues were evaluated for their inhibitory effects on HIV-induced cytopathogenicity in MT4 cells (Table 1, fig. 2).

Human T-lymphocyte MT4 cells (5×10^5 cells/ml) were suspended in fresh RPMI-1640 culture medium (Gibco) containing 10% (v/v) foetal calf serum (Gibco), 2 mM L-glutamine (Flow Laboratories), 20 mM Hepes buffer, 0.075% (w/v) NaHCO_3 (Flow Laboratories), 2.5 μ g/ml Fungizone (Squibb N.V., Brussels, Belgium) and 20 μ g/ml Geomycine (Essex N.V., Heist-o/d-Berg, Belgium), and infected with 400 CCID₅₀ - (Cell Culture Infective Dose-50) HIV per ml cell suspension. Then 100 μ l of the infected cell suspension was added to 100 μ l of an appropriate dilution of test compound in 200 μ l wells of a Flat Bottom Microtest III Plate (Falcon, Becton Dickinson, Oxnard, CA, USA) (i.e. 40 CCID₅₀ HIV/200 μ l well/ 5×10^4 cells), and further incubated at 37°C in a CO₂-controlled humidified atmosphere. After incubation for 5 days, viable cell counts were determined for both virus-infected cell cultures and non-infected cell cultures (which had been incubated with the same concentration of compounds as the virus-infected cells). The 50% effective dose (ED₅₀) and 50% cytotoxic dose (CD₅₀) were defined as the compound concentrations required to reduce by 50% the number of viable cells in the virus-infected and non-infected cell cultures, respectively.

TABLE 1 ANTI-RETROVIRUS EFFECTS OF 3'-FLUORO- AND 3'-AZIDO-SUBSTITUTED PYRIMIDINE 2',3'-DIDEOXYNUCLEOSIDE ANALOGUES

Compound	HIV-induced cyto-			MSV-induced cyto-		
	pathogenicity			pathogenicity		
	ED ₅₀ (μ M)	CD ₅₀ (μ M)	S.I.	ED ₅₀ (μ M)	CD ₅₀ (μ M)	S.I.
	1)	2)	4)	1)	3)	4)
ddUrd	210	>625	> 3	>400	>400	-
FddUrd	0.04	16	400	>400	>400	-
AzddUrd	0.36	244	677	15	>400	>27
ddEtUrd	>625	>625	-	>400	>400	-
FddEtUrd	330	>625	>1.9	>400	>400	-
AzddEtUrd	64	418	6.5	>200	>200	-
ddCyd	0.3	40	120	25	>400	>16
FddCyd	16	26	1.6	284	>400	> 1.2
AzddCyd	-	-	-	70	>400	> 5.7

1) 50% antiviral effective dose, required to affect a 50% reduction in the cytopathic effect of HIV for MT4 cells or transformation of C3H cells by MSV.

2) 50% cytotoxic dose required to reduce the number of viable cells in the untreated MT4 cell cultures by 50%.

3) The parameter followed here was an alteration of normal cell morphology.

4) Selectivity index or ratio of CD₅₀ to ED₅₀.

Marked differences were noted in the anti-HIV potencies of the 3'-substituted pyrimidine 2',3'-dideoxyribosides. With an ED₅₀ of 0.04 μ M, FddUrd was the most potent, and with an ED₅₀ of >64 μ M, ddUrd, ddEtUrd, FddEtUrd and AzddEtUrd were the least potent HIV inhibitors.

Introduction of a 3'-azido- or 3'-fluoro-group also increased the anti-HIV potency of ddUrd. With an ED₅₀ of 0.04 μ M FddUrd was 5000-fold and with an ED₅₀ of 0.36 μ M AzddUrd was 600-fold more effective against HIV replication than the parent compound ddUrd (ED₅₀: 210 μ M). Although FddUrd was 9-fold more effective against HIV than AzddUrd, it was also more cytostatic. This resulted in a selectivity index of 400 for FddUrd as compared to 677 for AzddUrd.

Among the ddCyd derivatives, 3'-fluoro-ddCyd was substantially less potent as an anti-HIV agent than ddCyd. The anti-retroviral effect of AzddCyd was not examined in the MT4 model. However, when evaluated in another T-lymphocyte cell line (ATH8), AzddCyd proved much less effective than ddCyd (H. Mitsuya et al, AIDS, Modern Concepts and Therapeutic Challenges (Broder, S., ed.), Marcel Dekker Inc., New York and Basel, pp. 303-333 (1987)).

ddUrd and the substituted pyrimidine 2',3'-dideoxynucleoside analogues were examined for their

inhibitory effect on the transformation of mouse embryo C3H fibroblasts by Moloney murine sarcoma virus (MSV).

C3H cells were seeded at 20,000 cells per ml into wells of Costar Tissue Culture Cluster plates (48 wells per plate) and grown to confluency. Cell cultures were then infected by 150 foci-forming units of MSV during 90 min, whereafter the culture medium was replaced by 1 ml fresh medium containing different concentrations of the test compound. After 6 days, the transformation of the cell cultures was examined microscopically.

It was surprisingly found that FddUrd was devoid of any anti-MSV activity ($ED_{50} > 400 \mu M$), and AzddUrd showed the highest activity (ED_{50} : $15 \mu M$). Further none of the ddEtUrd derivatives had any inhibitory effect on C3H cell transformation by MSV. Among the ddCyd analogues, ddCyd had some activity (ED_{50} : $25 \mu M$); the 3'-substituted ddCyd derivatives were less effective (see also table 1).

EXAMPLE 2

CYTOSTATIC AND ANTIMETABOLIC EFFECTS OF FddUrd AND 3'-SUBSTITUTED PYRIMIDINE 2',3'-DIDEOXYNUCLEOSIDE ANALOGUES

FddUrd and the other 3'-fluoro- and 3'-azido-substituted derivatives of ddUrd, ddEtUrd and ddCyd were examined for their cytostatic effects on B-lymphoblast (Raji/0), T-lymphoblast (Molt-4F) and T-lymphocyte (CEM, H9, MT4) cell lines.

All assays were performed in flat bottom Microtest III Plates (96 wells) (Falcon) as previously described by E. De Clercq et al (Mol. Pharmacol. 19, 321-330 (1981)). Briefly, the cells were suspended in growth medium and added to the microplate wells at a density of 7.5×10^4 cells/well ($200 \mu l$) in the presence of varying concentrations of the test compounds. The cells were then allowed to proliferate for 68-72 hours at $37^\circ C$ in a humidified, CO_2 -controlled atmosphere. At the end of the incubation period, the cells were counted with a Coulter Counter (Coulter Electronics Ltd., Harpenden Herts, U.K.). The 50% inhibitory dose (ID_{50}) was defined as the concentration of compound that reduced the number of cells by 50%.

Marked differences emerged, depending on the cell line; CEM cells were 70-fold more susceptible to the cytostatic action of FddUrd (ID_{50} : $14 \mu M$) than the other cell lines ($ID_{50} > 1000 \mu M$) (see table 2).

TABLE 2 CYTOSTATIC EFFECT OF 3'-SUBSTITUTED PYRIMIDINE
2',3'-DIDEOXYNUCLEOSIDE ANALOGUES AGAINST HUMAN B-
AND T-CELL LINES

Compound	ID ₅₀ ¹⁾ (μM)					
	Raji/0	Raji/TK ⁻	Molt/4F	CEM	H9	MT4
ddUrd	>1000	>1000	>1000	>1000	>1000	≥1000
FddUrd	>1000	>1000	≥1000	14	>1000	>1000
AzddUrd	≥1000	>1000	>1000	423	>1000	364
ddEtUrd	>1000	>1000	>1000	>1000	>1000	326
FddEtUrd	>1000	>1000	>1000	>1000	>1000	>1000
AzddEtUrd	>1000	>1000	>1000	>1000	>1000	160
ddCyd	23	28	7	6	243	≥1000
FddCyd	95	118	24	7	>1000	>1000
AzddCyd	-	-	-	-	-	-

1) 50% inhibitory dose required to reduce the cell number by 50%.

None of the 3'-substituted ddEtUrd derivatives, including ddEtUrd itself, were potent inhibitors of cell proliferation. Among the 2',3'-dideoxycytidine derivatives, ddCyd and 3'-FddCyd were more inhibitory to the proliferation of CEM and Molt/4F cells (ID₅₀: 6-24 μM) than Raji (ID₅₀: 23-95 μM), H9 and MT4 cells (243- >1000 μM).

EXAMPLE 3

IMPACT ON THE CELLULAR DNA SYNTHESIS

The impact of ddUrd and the 3'-fluoro- and 3'-azido-substituted pyrimidine 2',3'-dideoxynucleoside analogues on cellular DNA biosynthesis was evaluated by monitoring the effect of these compounds on the incorporation of radio-labelled (1',2'-³H)dUrd (specific radioactivity 27 Ci/mmol), (methyl-³H)dThd (specific radioactivity 480 Ci/mmol) and (5-³H)dCyd (specific radioactivity 20 Ci/mmol) into DNA of Molt/4F cells.

The incorporation of radiolabelled precursors into Molt/4F cellular DNA was measured in microtest III plates (96 wells) (Flacon) as previously described (J. Balzarini et al, Mol. Pharmacol. 23, 175-181 (1983)). Shortly, to each well of the microplate were added 10⁵ Molt/4F cells and 0.25 μCi of (1',2'-³H)dUrd, (methyl-³H)dThd or (5-³H)dCyd. The cells were allowed to proliferate for 20-24 hours at 37°C in a humidified CO₂-controlled atmosphere. At the end of this incubation period, the contents of the wells (200 μl) were brought onto 25-mm glass-fibre filters (type A/E, Gelman Instrument Company, Ann Arbor, Mich., USA) and

washed twice with cold phosphate-buffered saline, twice with cold 10% trichloroacetic acid (TCA), twice with cold 5% TCA, once with ethanol and once with ether. The filters were then assayed for radioactivity in a toluene-based scintillant.

ddUrd and ddEtUrd had no effect on the incorporation of (1',2'-³H)dUrd, (methyl-³H)dThd and (5-³H)-dCyd into Molt-4F cell DNA (Table 3).

TABLE 3

INHIBITORY ACTIVITY OF 3'-FLUORO- AND 3'-AZIDO-SUBSTITUTED PYRIMIDINE 2',3'-DIDEOXYNUCLEOSIDES ON THE INCORPORATION OF (1',2'-³H)dUrd, (methyl-³H)dThd AND (5-³H)dCyd INTO MOLT/4F CELL DNA

Compound	ID ₅₀ ¹⁾ (μM)		
	(1',2'- ³ H)dUrd	(methyl- ³ H)dThd	(5- ³ H)dCyd
	incorporation	incorporation	incorporation
ddUrd	>1000	>1000	>1000
FddUrd	16	>1000	>1000
AzddUrd	172	>1000	725
ddEtUrd	>1000	>1000	>1000
FddEtUrd	>1000	>1000	>1000
AzddEtUrd	>1000	>1000	>1000
ddCyd	32	753	>1000 (stimulation)
FddCyd	27.3	>1000	>1000 (stimulation)
AzddCyd	-	-	-

¹⁾ 50% inhibitory dose required to reduce the incorporation of the radiolabelled precursor by 50%.

FddUrd and AzddUrd inhibited (1',2'-³H)dUrd incorporation into DNA to a greater extent than (methyl-³H)dThd or (5-³H)dCyd incorporation. None of the ddEtUrd derivatives showed any effect on the incorporation of the radiolabelled DNA precursors. When ddCyd and FddCyd were evaluated for their inhibitory effects on (1',2'-³H)dUrd, (methyl-³H)dThd and (5-³H)dCyd incorporation, both compounds proved 25- to >40-fold more inhibitory to (1',2'-³H)dUrd incorporation than (methyl-³H)dThd incorporation. They significantly stimulated (5-³H)dCyd incorporation into DNA at the highest concentrations tested (200 μM and 1000 μM). The stimulatory effect of both ddCyd and FddCyd on (5-³H)dCyd incorporation into DNA amounted to 1.7-fold at 200 μM, and to 2-fold at 1000 μM, as compared to the control (data not shown).

EXAMPLE 4

Effect of FddUrd on the inhibition of tritium release from (5-³H) dUrd or (5-³H) dCyd in Molt 4F cells

The procedure to measure tritium release from (5-³H)dUrd or (5-³H)dCyd in intact cells has been described previously (J. Balzarini et al. Biochem. Biophys. Acta 785, 36-45 (1984)). Briefly, 10⁷ Molt 4F cells/ml were preincubated with an appropriate amount of test compound for 15 min at 37 °C. After this incubation period, radiolabelled (5-³H)dUrd or (5-³H)dCyd (100 µCi/ml; 0.1 µM) were added, and at various times (0, 15, 30, 45, 60 min), 100 µl of the reaction mixture was withdrawn, and mixed with 500 µl of a cold suspension of carbon black (100 mg/ml) in 5% TCA. After centrifugation at 1000 g for 10 min, the supernatants were analysed for radioactivity.

TABLE 4

INHIBITORY ACTIVITY OF 3'-FLUORO- AND 3'-AZIDO-SUBSTITUTED PYRIMIDINE 2',3'-DIDEOXYNUCLEOSIDES ON TRITIUM RELEASE FROM (5-³H)dUrd AND (5-³H)dCyd IN MOLT/4F CELLS

Compound	ID ₅₀ ¹⁾ (µM)	
	(5- ³ H)dUrd	(5- ³ H)dCyd
ddUrd	>250	>250
FddUrd	126	>250
AzddUrd	20	>250
ddEtUrd	>250	-
FddEtUrd	>250	-
AzddEtUrd	51	>250

1) 50% inhibitory dose required to reduce tritium release by 50%.

None of the compounds evaluated affected tritium release from (5-³H)dCyd even at a concentration of 250 µM (Table 4). Among the 3'-substituted ddUrd derivatives, AzddUrd, and to a lesser extent FddUrd inhibited tritium release from (5-³H)dUrd (ID₅₀: 20 and 126 µM, respectively). AzddEtUrd was the only ddEtUrd derivative that inhibited tritium release from (5-³H)dUrd (ID₅₀: 51 µM) (Table 4.)

EXAMPLE 5

Affinity of Molt/4F dThd kinase and dCyd kinase for FddUrd

ddUrd and the 3'-fluoro- and 3'-azido-substituted pyrimidine 2',3'-dideoxynucleoside analogues were evaluated for their inhibitory effect against dThd and dCyd kinase.

dThd kinase and dCyd kinase were prepared from exponentially growing Molt 4F cells, which were first washed (2x) with phosphate-buffered saline at 4° and then homogenized by sonication. The suspensions were clarified by centrifugation at 25,000-27,000 x g for 30 min. The 30-70% (NH₄)₂SO₄ precipitate of the supernatant was resuspended in buffer containing 10 uM potassium phosphate, pH 7.5, 10 mM β-mercaptoethanol and 0.1 M KCl, dialyzed against the same buffer, and used for the dThd kinase and the dCyd kinase assays. In these experiments, (2-¹⁴C)dThd and (U-¹⁴C)dCyd served as the radiolabelled substrates. The apparent K_m and K_i values were derived from Lineweaver-Burk plots, using a linear regression analysis program. The assay procedures have been described in detail (J. Balzarini et al, (1983) and J. Balzarini et al, Biochem. Pharmacol. 31, 3673-3682 (1982)).

TABLE 5 INHIBITION OF MOLT/4F dThd KINASE AND dCyd KINASE BY 3'-SUBSTITUTED PYRIMIDINE 2',3'-DIDEOXYNUCLEOSIDE ANALOGUES

Compound	Enzyme	K _i	K _i /K _m ¹⁾	Type of inhibition
ddUrd	dThd kinase	>1000	>250	-
FddUrd	dThd kinase	833	171	competitive
AzddUrd	dThd kinase	224	71	competitive
ddEtUrd	dThd kinase	>750	>200	-
FddEtUrd	dThd kinase	266	72	competitive
AzddEtUrd	dThd kinase	175	28	competitive
ddCyd	dCyd kinase	1965	404	competitive
FddCyd	dCyd kinase	360	63	competitive
AzddCyd	dCyd kinase	354	58	competitive

¹⁾ K_m values obtained in the individual experiments ranged from 2.5 to 6.6 μM for dThd kinase and 5.2 to 6.4 μM for dCyd kinase.

The K_i and K_i/K_m values of Molt/4F dThd kinase for ddUrd, ddEtUrd, and their 3'-fluoro- and 3'-azido-substituted derivatives are presented in Table 5. The unsubstituted 2',3'-dideoxynucleosides (i.e. ddUrd and ddEtUrd) had a substantially lower affinity for dThd kinase than their 3'-fluoro- and 3'-azido-substituted counterparts. Moreover, the 3'-azido substituent conferred a greater increase in affinity of the 2',3'-dideoxynucleoside for dThd kinase than did the 3'-fluoro substituent. The K_i/K_m values for FddUrd and AzddUrd being 171 and 71, respectively. For ddUrd and ddEtUrd no inhibitory activity against dThd kinase could be detected, even when evaluated at a concentration as high as 750 μM or 1000 μM (Table 5). For those ddUrd and ddEtUrd analogues that were active as inhibitors of dThd kinase, the type of inhibition appeared to be competitive with respect to dThd. Lineweaver-Burk plots are shown for FddUrd, and AzddUrd (Fig. 3).

The K_i , K_m values of Molt 4F dCyd kinase for ddCyd, FddCyd and AzddCyd are also presented in Table 5. The K_m value of the enzyme for dCyd ranged from 5.2 μ M to 6.4 μ M. FddCyd and AzddCyd had a relatively low affinity for the enzyme (K_i , $K_m \sim 60$), and for the parent nucleoside ddCyd the K_i , K_m value was even higher (Table 5). In all cases, inhibition was competitive with respect to radiolabelled dCyd as the competing substrate. Lineweaver-Burk plots are shown for FddCyd and AzddCyd in Fig. 4.

EXAMPLE 6

Kinetic properties of FddUrd for bacterial thymidine phosphorylase

Phosphorolytic cleavage of the test compounds was assessed using purified bacterial dThd phosphorylase (Sigma Chemical Co., St. Louis, MO) with the aid of a Beckman spectrophotometer UV Vis 5200; the absorbance change during the enzyme reaction was continuously monitored at 25°C at the wavelength where the difference of absorption between the nucleoside and its free base was maximal. The reaction mixtures for all the assays consisted of enzyme and non-saturating concentrations of nucleoside in 0.1 M sodium phosphate buffer, pH 7.0. The initial velocity of phosphorolysis was expressed as nmol.min:enzyme unit.

The 3'-fluoro- and 3'-azido-substituted, as well as the unsubstituted, ddUrd and ddEtUrd nucleoside analogues were evaluated for their susceptibility to phosphorolysis by purified bacterial dThd phosphorylase. As compared to the 2'-deoxyribosides dUrd and dEtUrd, the 2',3'-dideoxyribosides showed a 1000- to 2000-fold decrease in the velocity of phosphorolysis by dThd phosphorylase (Table 6) (see also J. Balzarini et al, Mol. Pharmacol. 32, 162-167 (1987)). For ddEtUrd no phosphorolytic cleavage whatsoever was detectable, and this was also the case for FddEtUrd and AzddEtUrd. Introduction of a fluorine at C-3' of ddUrd further decreased the affinity for dThd phosphorylase. Introduction of an azido group at C-3' of ddUrd made the compound even totally resistant to hydrolysis by dThd phosphorylase. Similarly, ddCyd, FddCyd and AzddCyd proved totally resistant to the action of dThd phosphorylase.

**TABLE 6 KINETIC PROPERTIES OF 3'-SUBSTITUTED PYRIMIDINE
2',3'-DIDEOXYNUCLEOSIDE ANALOGUES FOR BACTERIAL
THYMIDINE PHOSPHORYLASE**

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Compound	Initial velocity of phosphorolysis (nmol/min/unit)
dUrd	361
ddUrd	0.270
AzddUrd	N.D. ¹⁾
FddUrd	0.018
dEtUrd	0.893
ddEtUrd	N.D.
AzddEtUrd	N.D.
FddEtUrd	N.D.
dCyd	N.D.
ddCyd	N.D.
AzddCyd	N.D.
FddCyd	N.D.

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¹⁾ Phosphorolysis not detectable with 25 units of enzyme.

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All examples demonstrate that FddUrd, whose synthesis and biological activity has not been previously revealed, is a potent and selective anti-HIV agent and should be considered as a novel candidate compound in the treatment of retrovirus infections, for instance AIDS and AIDS-related diseases, including hepatitis B.

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The introduction of a fluorine at C-3' of a pyrimidine 2',3'-dideoxynucleoside, does not inherently lead to an increase of anti-HIV activity. 3'-fluoro-substitution of ddUrd leads to an increase of anti-HIV activity, but in the case of ddCyd leads to a decrease in anti-retrovirus activity. Further all 3'-substituted ddEtUrd derivatives show little, if any, anti-retrovirus activity (see table 1, 5).

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The differential effects of 3'-fluoro and 3'-azido substitution on the biological activity of ddUrd and ddEtUrd on the one hand, and ddCyd on the other, may be related to differences in the affinity of these compounds for the kinases involved in their phosphorylation, as well as differences in the affinity of the resulting 5'-triphosphates for the reverse transcriptase.

Claims

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1. 3'-fluoro-2',3'-dideoxyuridine.
2. A therapeutic composition for use in the treatment of retroviral diseases which comprises 3'-fluoro-2',3'-dideoxyuridine as an active ingredient.

3. A therapeutic composition as claimed in claim 2, comprising 3'-fluoro-2',3'-dideoxyuridine in a concentration ranging from 0.1 - 100% by weight.

4. A therapeutic composition as claimed in claim 3, having the form which is selected from the group consisting of powders, suspensions, solutions, sprays, emulsions, unguents and creams.

5. A therapeutic composition for use in the treatment of AIDS or AIDS-related diseases including hepatitis B, which comprises 3'-fluoro-2',3'-dideoxyuridine as an active ingredient.

6. A therapeutic composition as claimed in claim 5, comprising 3'-fluoro-2',3'-dideoxyuridine in a concentration ranging from 0.1 - 100% by weight.

7. A therapeutic composition as claimed in claim 6, having the form which is selected from the group consisting of powders, suspensions, solutions, sprays, emulsions, unguents and creams.

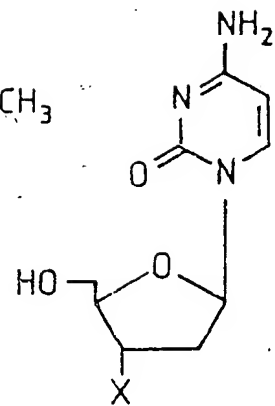
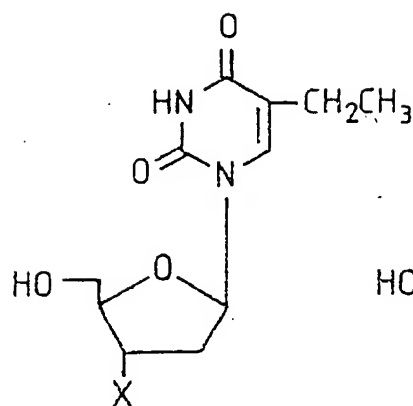
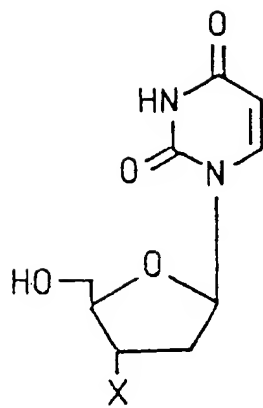
8. A method for the treatment of a retroviral disease which comprises administering 3'-fluoro-2',3'-dideoxyuridine to a patient suffering from the retroviral disease.

9. A method for the treatment of AIDS or AIDS-related diseases which comprises administering 3'-fluoro-2',3'-dideoxyuridine to a patient suffering from AIDS or AIDS-related diseases, including hepatitis B.

10. Use of 3'-fluoro-2',3'-dideoxyuridine for preparing a therapeutic composition against a retroviral disease.

11. Use of 3'-fluoro-2',3'-dideoxyuridine for preparing a therapeutic composition against AIDS or AIDS-related diseases, including hepatitis B.

FIG. 1



X

H

ddUrd

ddEtUrd

ddCyd

F

FddUrd

FddEtUrd

FddCyd

N₃

AzddUrd

AzddEtUrd

AzddCyd

FIG. 2

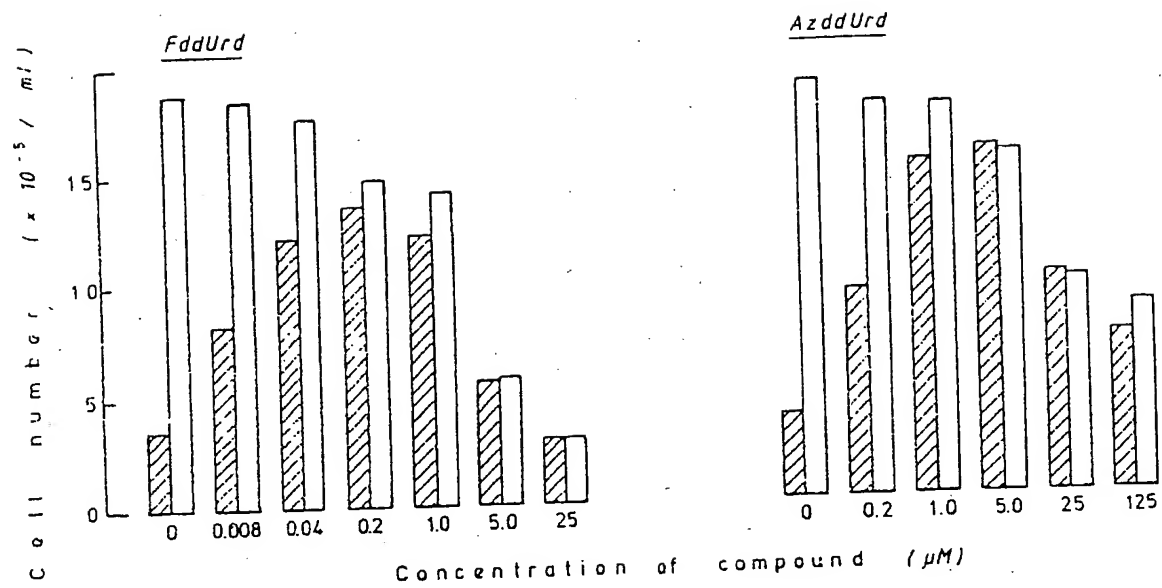


FIG. 3

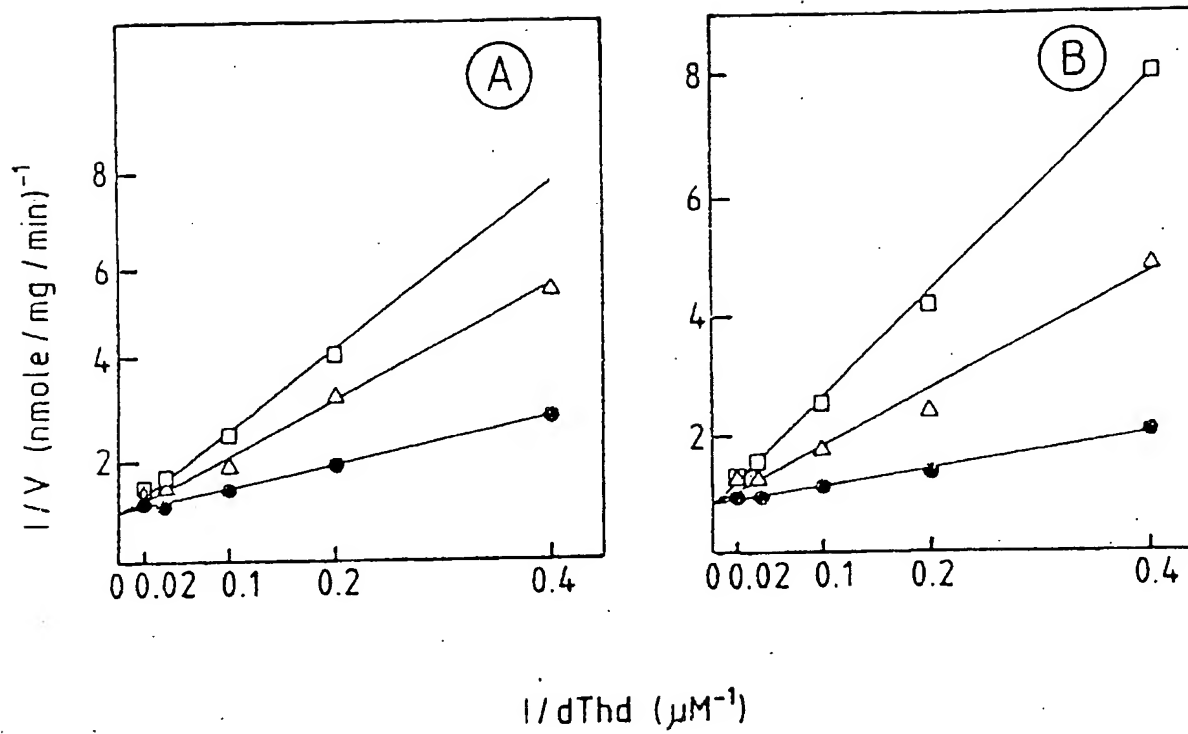
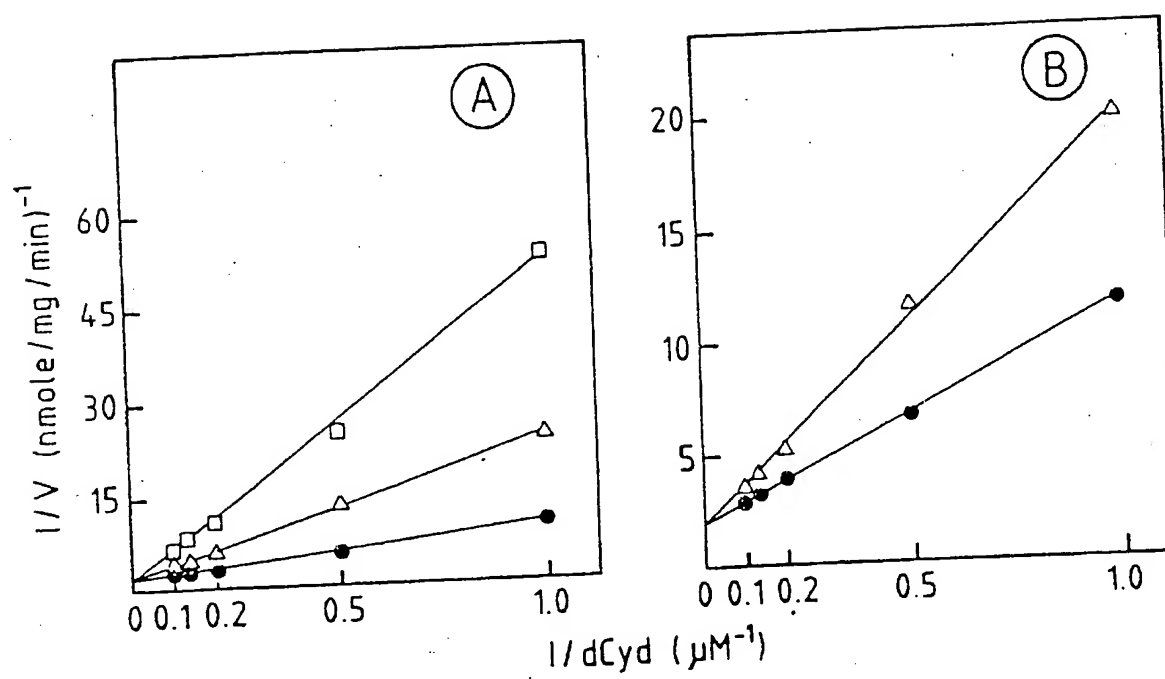


FIG. 4





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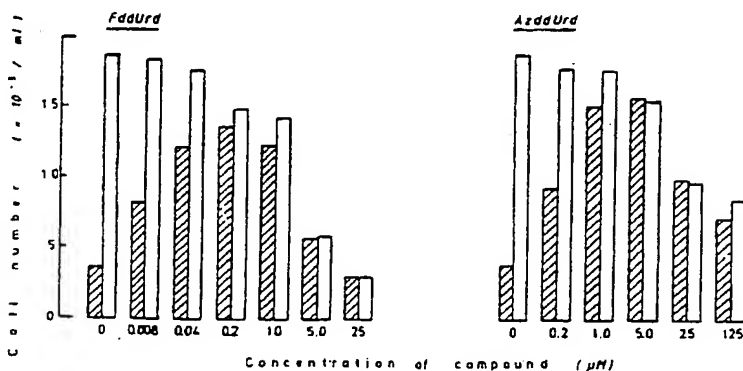
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3'-Fluoro-2',3'-dideoxyuridine, and its therapeutic application.

The novel compound 3'-fluoro-2',3'-dideoxyuridine has an antiretroviral effect against human immunodeficiency virus (HIV) and is a potential an-

tiretroviral agent for treating AIDS and related viral diseases, including hepatitis B.

FIG.2



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which under Rule 45 of the European Patent Convention
shall be considered, for the purposes of subsequent
proceedings, as the European search report

Application number

EP 88 11 7249

DOCUMENTS CONSIDERED TO BE RELEVANT			
Category	Citation of document with indication, where appropriate, of relevant passages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. Cl.4)
X	US-A-3 775 397 (DEUTSCHE AKADEMIE DER WISSENSCHAFTEN ZU BERLIN) * The whole document *	1	C 07 H 19/073 A 61 K 31/70
X	THE JOURNAL OF MEDICINAL CHEMISTRY, vol. 30, no. 8, August 1987, pages 1270-1278, American Chemical Society; P. HERDEWIJN et al.: "3'-Substituted 2',3'-dideoxynucleoside analogues as potential anti-HIV (HTLV-III/ LAV) agents" * Abstract; compounds 3d,3g,3i,3j *	1,2, 5,10, 11	
X,P	EP-A-0 254 268 (DEUTSCHE AKADEMIE DER WISSENSCHAFTEN DER DDR) ./.		TECHNICAL FIELDS SEARCHED (Int. Cl.4) C 07 H 19/00 A 61 K 31/00
INCOMPLETE SEARCH The Search Division considers that the present European patent application does not comply with the provisions of the European Patent Convention to such an extent that it is not possible to carry out a meaningful search into the state of the art on the basis of some of the claims. Claims searched completely: 1-7,10,11 Claims searched incompletely: Claims not searched: 8,9 Reason for the limitation of the search: Method for treatment of the human or animal body by surgery or therapy (see Art. 52(4) of the European Patent Convention).			
Place of search The Hague		Date of completion of the search 15-06-1990	Examiner SCOTT
CATEGORY OF CITED DOCUMENTS X : particularly relevant if taken alone Y : particularly relevant if combined with another document of the same category A : technological background O : non-written disclosure P : intermediate document T : theory or principle underlying the invention E : earlier patent document, but published on, or after the filing date D : document cited in the application L : document cited for other reasons & : member of the same patent family, corresponding document			



DOCUMENTS CONSIDERED TO BE RELEVANT			CLASSIFICATION OF THE APPLICATION (Int. Cl. 4)
Category	Citation of document with indication, where appropriate, of relevant passages	Relevant to claim	
	<p>* Abstract; page 3, line 1 - page 4, line 35; claims 1,9-13 *</p> <p>--</p>	1-7, 10,11	
X	<p>CHEMICAL ABSTRACTS, vol. 80, no. 9, March 4, 1974, page 392, abstract 48302d, Columbus, Ohio, US; G. KOWOLLIK et al.: "Nucleosides of fluorinated sugars. XII. Synthesis of 1-(2,3-dideoxy-3-fluoro-beta-D-ribofuranosyl)pyrimidines" & J. PRAKT. CHEM. 1973, 315(5), 895-900</p> <p>* Abstract *</p> <p>--</p>	1	TECHNICAL FIELDS SEARCHED (Int. Cl. 4)
X	<p>CHEMICAL ABSTRACTS, vol. 78, no. 21, May 28, 1973, page 414, abstract 136580g, Columbus, Ohio, US; D. CECH et al.: "Reaction of uracil and its nucleosides with elemental fluorine" & J. PRAKT. CHEM. 1973, 315(1), 149-54</p> <p>* Abstract *</p> <p>-----</p>	1	

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